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Specific ultra-violet absorbance as an indicator of mercury sources in an Adirondack River basin

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Abstract The Adirondack region of New York has been identified as a hot spot where high methylmercury concentrations are found in surface waters and biota, yet mercury (Hg) concentrations vary widely in this region. We collected stream and groundwater samples for Hg and organic carbon analyses across the upper Hudson River, a 493 km² basin in the central Adirondacks to evaluate and model the sources of variation in filtered total Hg (FTHg) concentrations. Variability in FTHg concentrations during the growing seasons (May–Oct) of 2007–2009 in Fishing Brook, a 66-km² sub-basin, was better explained by specific ultra-violet absorbance at 254 nm (SUVA₂₅₄), a measure of organic carbon aromaticity, than by dissolved organic carbon (DOC) concentrations, a commonly used Hg

indicator. SUVA₂₅₄ was a stronger predictor of FTHg concentrations during the growing season than during the dormant season. Multiple linear regression models that included SUVA₂₅₄ values and DOC concentrations could explain 75 % of the variation in FTHg concentrations on an annual basis and 84 % during the growing season. A multiple linear regression landscape modeling approach applied to 27 synoptic sites across the upper Hudson basin found that higher SUVA₂₅₄ values are associated with gentler slopes, and greater riparian area, and lower SUVA₂₅₄ values are associated with an increasing influence of open water. We hypothesize that the strong Hg–SUVA₂₅₄ relation in this basin reflects distinct patterns of FTHg and SUVA₂₅₄ that are characteristic of source areas that control the mobilization of Hg to surface waters, and that the seasonal influence of these source areas varies in this heterogeneous basin landscape.

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Introduction

Organic matter regulates mercury (Hg) cycling in the environment through numerous mechanisms including: ligand binding, effects on dissolution and precipitation, by serving as a microbial substrate, through its central role in redox processes, and as an indicator of hydrologic conditions (Gerbig et al. 2012). The strong

association of Hg and organic matter in surface waters has long been recognized (Lindberg and Harriss 1974; Mierle and Ingram 1991), and numerous studies of Hg in streams and lakes of mid- and northern latitudes of the northern hemisphere have generally shown strong correlations between dissolved Hg concentrations and dissolved organic carbon (DOC) concentrations, and between particulate Hg concentrations and particulate organic carbon concentrations (POC) (Grigal 2002; Yin and Balogh 2002; Dennis et al. 2005; Brigham et al. 2009; Riscassi and Scanlon 2011). The association of Hg with organic matter is sufficiently strong in many surface waters, that measures such as DOC or absorbance at 254 nm (UV_{254}) have been suggested as surrogates for Hg concentrations, because Hg analyses are expensive and may be unstable in automated water samplers (Dittman et al. 2009).

Interactions between Hg and organic matter are numerous and diverse, reflecting the complexity of natural organic matter (Ravichandran 2004; Gerbig et al. 2012). Strong binding of Hg by sulfur-bearing thiol groups has been shown, and is believed to be a strong driver of Hg association with dissolved organic matter (DOM) (Benoit et al. 2001; Skjellberg 2008). Hg may also be bound to organic matter as HgS nanoparticles that pass through 0.45 μm pore size filters, but this form may not be common in surface waters unimpacted by mining activities (Gerbig et al. 2011).

In natural waters, measures that reflect in part, the relative aromaticity of DOM such as the hydrophobic acid (HPOA) concentration or UV_{254} , are often more strongly related to dissolved Hg than a general measure such as DOC or TOC (Shanley et al. 2008; Dittman et al. 2010; Riscassi and Scanlon 2011). Among several possible reasons for this association is that most reduced sulfur (and therefore, most of the strong Hg binding sites) is typically found within the HPOA fraction of DOM (Ravichandran 2004), which is more aromatic than the non-HPOA fraction. Additionally, the more aromatic fraction of organic matter may better stabilize Hg–S nanoparticles (Ravichandran et al. 1999; Gerbig et al. 2011). Thus, the association of Hg with the aromatic fraction of organic matter may be rooted in the complex chemistry of Hg interactions with natural organic matter. In addition to geochemistry-based reasons why Hg is more strongly associated with aromatic organic matter, this association may also originate through mobilization to

surface waters and mixing of Hg and DOM from principal runoff source areas such as humic-rich shallow soils (Dittman et al. 2010) as well as riparian wetland soils (Ågren et al. 2008; Mierle and Ingram 1991), two locations where preferential storage of Hg has been identified (Grigal 2003). In summary, Hg–DOM chemistry, storage, and hydrology are interrelated in complex ways that are the subject of ongoing research; thus, determining the primacy of any single explanation for an association that is evident at the basin scale may be difficult.

Although dissolved Hg and DOC concentrations are often strongly correlated in many surface waters with r^2 values of 0.7–0.9 (Grigal 2002; Shanley et al. 2008; Dittman et al. 2010; Riscassi and Scanlon 2011), not all waters show such strong a strong association. For example, Hg and DOC concentrations are weakly related in the Everglades where a diversity of organic matter sources and complex biogeochemical processes are observed (Aiken et al. 2011). Similarly, in surface waters strongly influenced by urban land use, diverse sources of Hg and DOC such as waste water treatment facilities often preclude a simple linear relation between dissolved Hg and DOC concentrations (Brigham et al. 2009).

The Adirondack region of New York is a largely forested and mountainous area with abundant lakes, streams, and wetlands. Surface waters in this region have been the frequent focus of investigations because of high sensitivity to acidification from acid deposition, and also due to high levels of Hg measured in surface waters and biota (Jenkins et al. 2005; Evers et al. 2007). Studies in Adirondack surface waters similar to those in nearby New England and southeastern Canada have shown a strong association of dissolved Hg and DOC concentrations (Driscoll et al. 1995; Selvendiran et al. 2009; Dittman et al. 2010). Many previously published investigations, however, have focused on small catchments or small lakes with drainage areas of only a few km^2 , and relatively homogeneous forested landscapes. Here, we describe results of a study in the 493 km^2 upper Hudson River basin, which drains a largely forested area of diverse topography that includes mountainous terrain and riparian floodplains dominated by wetlands as well as numerous lakes and ponds interconnected by streams and rivers. Recently published studies in this basin have shown a relatively weak correlation between total Hg (THg) and DOC concentrations that varies

seasonally and with flow (Bushey et al. 2008; Schelker et al. 2011; Burns et al. 2012), though some data in this basin indicate a strong Hg–DOC relation (Selvendiran et al. 2009; Dittman et al. 2010). We argue that the diverse landscape in this basin results in a range of organic matter sources and sinks that greatly affect the cycling of Hg and preclude the strong correlation between dissolved Hg and DOC concentrations often observed in smaller more homogeneous basins in this region. Here, we build on previous work that indicated specific ultra-violet absorbance at 254 nm ($SUVA_{254}$) was a stronger predictor of variation in THg concentrations than were DOC concentrations during the summer across the upper Hudson basin (Burns et al. 2012). We hypothesize that $SUVA_{254}$ is a potentially powerful predictor of dissolved Hg concentrations because this measure reflects the source areas and landscape locations where DOC and Hg interactions occur as well as the relative aromaticity and ability of the DOC to bind and transport Hg. Furthermore, we use a multiple regression spatial modeling approach to examine the relative influence of DOC concentrations

and $SUVA_{254}$ values on Hg concentrations, and use a similar approach to explore the landscape elements that most greatly affect variation in $SUVA_{254}$ values. The results reported here may differ from those in small catchments because the Hg cycle in larger heterogeneous basins is affected by a diverse array of landscape types and varied sources and sinks of organic matter. Studies of Hg cycling in larger basins are important because these surface waters are more likely to have complex food webs that extend to larger carnivorous fish, which tend to have the highest Hg levels, and therefore pose the greatest human health risk.

Study site

The upper Hudson River is defined here as the 492.8 km² basin upstream of the U.S. Geological Survey gage near Newcomb, NY in the Adirondack State Park (Fig. 1). Fishing Brook at County Line Flow is a 65.6 km² sub-basin that defines the

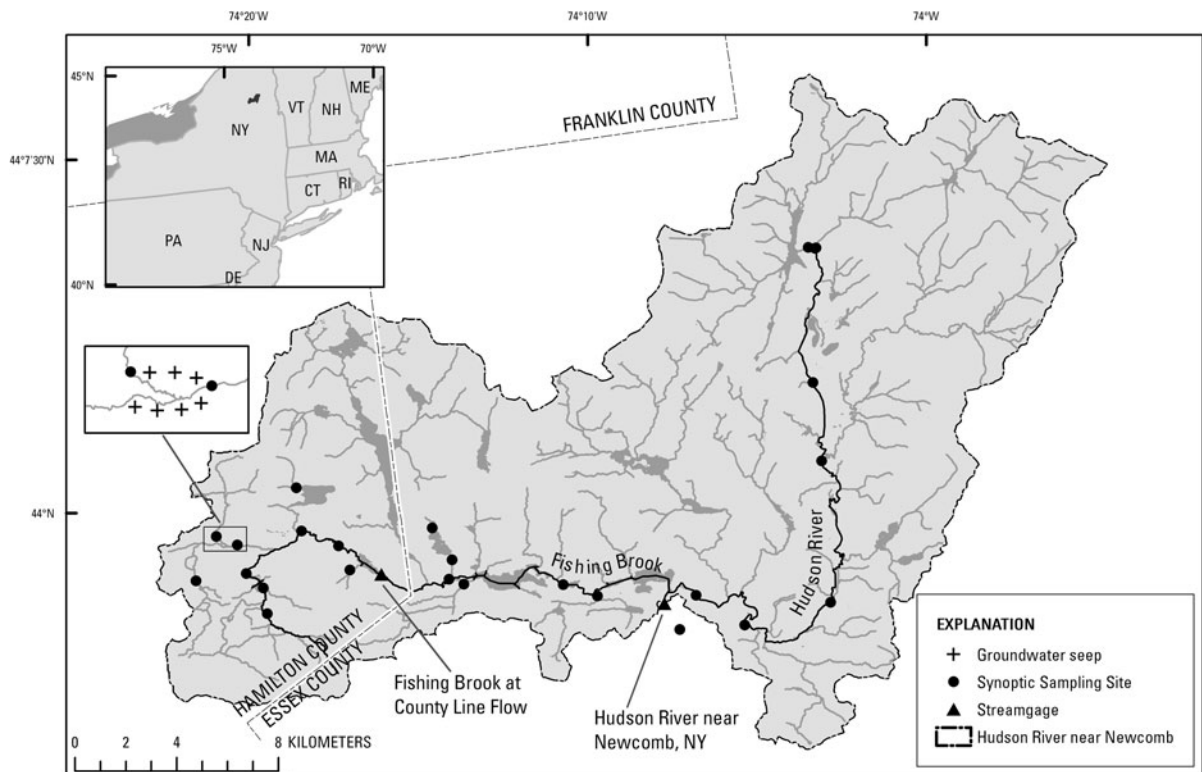


Fig. 1 Map showing gaged stream sampling sites, Fishing Brook at County Line Flow and the Hudson River near Newcomb as well as 25 additional synoptic sampling sites discussed in paper. Locations of seven groundwater seeps shown on *inset* map

westernmost boundary of the upper Hudson basin (Fig. 1). The upper Hudson basin is underlain by metamorphic bedrock of middle Proterozoic age (Chiarenzelli and McLelland 1993). There are distinct steep (mean slope = 30 %) upland areas in the northernmost and westernmost parts of the basin overlain by coarse rocky till with exposed bedrock, and riparian floodplains that are dominant at lower elevations in the basin overlain by sandy outwash and alluvium with intermixed discontinuous peat (NRCS 2006, 2010). Much, though not all of this riparian area is classified as wetland; 6.7 % of the basin is wetland area (LaPoint et al. 2004; Dahl et al. 2009). The basin also has many ponds and lakes; 3.6 % of the basin is classified as open water (excluding wetland area; 1:24,000 National Hydrography Dataset; <http://nhd.usgs.gov/data.html>; accessed 17 Jan 2012). The basin is largely forested (92 %) (1992 National Land Cover Dataset; <http://landcover.usgs.gov/natl/landcover.php>; accessed 17 Jan 2012), with <1 % developed land. Elevation ranges from 423 to 1624 m. The basin is in the southernmost part of the Eastern forest—boreal transition ecoregion (Olson et al. 2001), and is dominated by northern hardwood forest below 1,000 m and spruce-fir at higher elevations and in wet areas adjacent to surface waters.

Methods

Two principal data sets were analyzed in this paper: (1) Hg and organic carbon fraction concentrations in 43 stream samples collected from Fishing Brook at County Line Flow approximately every 3 weeks for 33-months from Jan 2007 to Sept 2009, and (2) SUVA₂₅₄ values in 54 synoptic stream samples collected at 27 catchments scattered across the upper Hudson basin during spring (May) and summer (July and Aug) of 2006 and 2008. Data from the Fishing Brook gage were analyzed in sum and by season (dormant Nov 1–Apr 30, growing May 1–Oct 31) because seasonal patterns were observed in past analyses at this site (Schelker et al. 2011). Additionally, results from 22 samples collected at 7 groundwater seeps located in the largest riparian wetland within Fishing Brook (Fig. 1) are also discussed. Sampling, analytical, and data analysis methods for these samples are described below.

Field sampling

Fishing Brook was sampled at the outlet of County Line Flow, a small shallow impoundment just upstream of the USGS stream gage that recorded stage at 15-min intervals for conversion to streamflow through a rating curve (Rantz et al 1982). All stream samples were collected by wading to the center point of flow or reaching to the center of flow when channels were narrow or to sample groundwater seeps. Each bottle was rinsed three times with stream water before filling.

A trace metal sampling technique similar to EPA Method 1669 (US EPA 1996) was used for Hg that employed wrist-length nitrile (use of name brands in this paper is for identification purposes only and does not represent an endorsement by the U.S. government) gloves over shoulder-length polyethylene gloves. Samples were filtered through 0.7- μ m pore size, prebaked (550 °C) quartz-fiber filters, acidified to approximately 1 % with 6 N HCl, and stored in the dark until analysis. The filters were placed in sterile petri dishes and stored frozen on dry ice until analysis. Additional details about Hg sampling methods are provided in Lewis and Brigham (2004) and QA/QC details are described in Schelker et al. (2011) and Burns et al. (2012).

Samples for DOC and UV₂₅₄ were pressure-filtered through baked glass fiber 0.7- μ m filters into 150 mL amber glass bottles and stored at 4 °C until analysis. The filter was also stored at 4 °C for later analysis of POC. Samples for HPOA analysis were passed through a 0.45- μ m capsule filter into 1 L amber glass bottles and similarly stored at 4 °C until analysis.

Laboratory analyses

Filtered total Hg (FTHg) and particulate total Hg (PTHg) were analyzed at the USGS Mercury Research Laboratory in Middleton, Wisconsin. Analysis of Hg was by cold vapor atomic fluorescence spectrometry according to the method described by Olson and DeWild (1999). THg is reported as the sum of FTHg and PTHg.

DOC concentrations, UV₂₅₄ measurements, and DOC fractionation analyses were performed at the USGS Organic Carbon Laboratory in Boulder, Colorado. Dissolved organic carbon was analyzed on an OI Analytical Model 700 TOC analyzer (Aiken et al.

1992). UV_{254} measurements (absorbance at $\lambda = 254$ nm) were made with a Hewlett-Packard Photodiode Array Spectrophotometer (model 8453) using a 10 mm quartz cell. UV_{254} data were corrected for the presence of iron (Fe) by subtracting the absorbance resulting from Fe(III) from the total absorbance value using an extinction coefficient of 0.0687; note that this coefficient is less than the value of 0.081–0.085 reported by Weishaar et al. (2003), and is based on more recent experimental data (B. Poulin personal communication). Analysis of Fe was done at the USGS National Water Quality Laboratory by Inductively-Coupled Plasma Emission Spectroscopy (USGS Method 01046, <https://www.nemi.gov>; accessed 17 Jan 2012). $SUVA_{254}$ values were obtained by dividing the corrected UV_{254} value by the DOC concentration (mg C/L) and are reported in units of L/mg C/m. The HPOA fraction was obtained following established protocols (Aiken et al. 1992; Spencer et al. 2010). In brief, samples were acidified to pH 2 with HCl and passed through a column of XAD-8 resin. The HPOA fraction was retained on the XAD-8 resin and then back eluted with 0.1 M NaOH. The percent HPOA was determined by dividing the HPOA concentration by that of DOC. POC was analyzed at the USGS National Water Quality Laboratory by high temperature combustion according to EPA Method 440.0 (<https://www.nemi.gov>; accessed 17 Jan 2012).

Landscape metrics

The landscape metrics explored for inclusion in multiple regression models that best predict $SUVA_{254}$ values across the 27 synoptic sites are the same as those discussed previously by Burns et al. (2012) for modeling THg and MeHg in the same basin. All of these metrics are described in Supplemental Table S-1, but only those that appeared in the best models are described here.

Landscape metrics were compiled for each sub-basin by geographic information system (GIS) analyses using Arc GIS software. The mean slope and mean topographic wetness index (TWI) were calculated for each sub-basin by terrain analysis of the 10 m National Elevation Dataset (<http://seamless.usgs.gov>, accessed 17 Jan 2012) using the System for Automated Geoscientific Analyses (SAGA, <http://www.saga-gis.org/en/index.html>, accessed 17 Jan 2012; Cimmery 2007). The TWI, $\ln(a/\tan \beta)$, was first

defined by Beven and Kirkby (1979), in which a is accumulated area per unit contour length and β is the local slope angle. Riparian area (expressed as proportion of total basin area and termed ripct) was calculated in SAGA by including all cells contributing to a given stream cell within an elevation <0.65 m above the stream cell. Wetland land cover was derived from the Adirondack Park Agency coverage, in which wetlands were delineated from 1:40,000 color infrared aerial photographs according to the approach used in the National Wetlands Inventory (Dahl et al. 2009; LaPoint et al. 2004).

Basin percent open water area is a commonly applied metric in studies of spatial controls on Hg cycling (Hurley et al. 1995; Shanley et al. 2005), and was explored here for inclusion in multiple regression models. An alternative to this metric, the open water index (OWI) that accounts for the proportion of basin drainage area that transits each open water body >0.025 km² was also calculated (described in greater detail by Burns et al. 2012). The remaining small open water bodies (<0.025 km² each) were compiled into a separate small open water body surface area metric for each sub-basin.

Statistical analyses

Multiple linear regression models were developed (SigmaStat software) to explore how well DOC concentrations and $SUVA_{254}$ values together explained variation in FTHg concentrations at Fishing Brook at County Line Flow. A forward stepwise method was applied, and both independent variables were significant ($p \leq 0.05$, with one noted exception) for all models discussed in the text. Additional multiple linear regression models were explored to determine which combination of the landscape metrics shown in Supplemental Table S-1 could best explain the variation in $SUVA_{254}$ values across the 27 synoptic sites in the upper Hudson basin. Models were explored for every possible combination up to eight variables. A sequence of steps was then followed to select the “best” model. First, any models in which the p value of one of the independent variables was >0.05 were rejected. Second, any models that included a variance inflation factor (VIF) >4.0 , a conservative level (O’Brien 2007), were rejected. The VIF is a measure of the proportion of variance accounted for by a given variable relative to the other independent variables in

the model, and thus can help avoid including variables with high multi-collinearity, a common occurrence for landscape metrics. In general, any model with more than four predictive variables was rejected based on either $p > 0.05$ or $VIF > 4.0$. Any models remaining after these two steps were further evaluated by selecting those with the highest R^2 values for each incremental number of independent variables, and then applying Akaike's Information Criteria to select the best model with the least number of independent variables.

Results and discussion

Seasonal variation in Hg and organic carbon measures

THg concentrations ranged from 1.3 to 3.9 ng/L (mean = 2.4 ng/L) in Fishing Brook during the study period of Jan 2007–Sept 2009 (Fig. 2a). The closeness of the symbols for THg and FTHg in Fig. 1a indicate that the THg fraction is dominated by filtered Hg. Mean FTHg/THg was 83.2 % and never less than 65 % in any sample collected. Similarly, the TOC and DOC symbols are quite close in Fig. 2a, and mean DOC/TOC was 94.4 % in Fishing Brook. Due to the dominance of the dissolved fractions for both THg and TOC, the analysis in this paper focuses on interpreting variation in DOC and FTHg concentrations.

Clear seasonal variation is apparent in the annual patterns of DOC and FTHg concentrations. For example, the highest annual DOC values of 9–12 mg/L were observed during summer ($p < 0.001$), when streamflow is typically lowest for the year. This type of seasonal pattern in DOC concentrations is commonly observed in watersheds dominated by near-stream wetlands (Schiff et al. 1998; Ågren et al. 2008), presumably due to: (1) waning influence of steeper upland areas with shallow soils during the driest seasonal conditions, and (2) increased influence from shallow groundwater in near-stream riparian areas with high DOC concentrations, where water tables remain relatively high in summer (Vidon and Hill 2004). In contrast, the greatest DOC and TOC concentrations in upland watersheds that lack significant near-stream wetlands or peat commonly occur

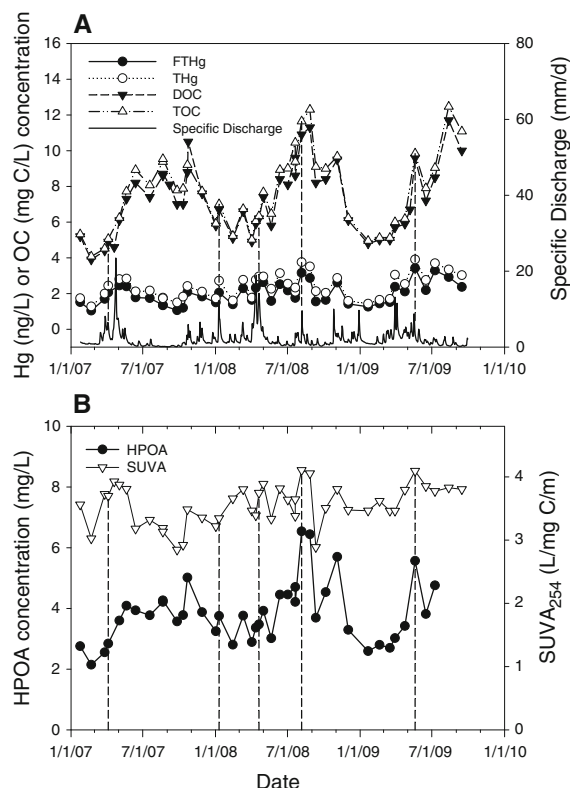


Fig. 2 Measures of Hg and OC in Fishing Brook at County Line Flow for samples collected during 2007–2009: **a** THg, FTHg, DOC, and TOC concentrations, **b** HPOA concentrations and SUVA₂₅₄ values. The five vertical dashed lines indicate samples collected at or near hydrograph peaks and are discussed in text

during snowmelt and rain storms, driven by the flushing of organic carbon from shallow soils (Boyer et al. 1997; Hinton et al. 1997). Such watersheds often have low baseflow DOC concentrations of 1–5 mg/L with higher values during leaf fall (Mulholland 2003). Event-driven DOC flushing patterns are not readily apparent in Fishing Brook (Schelker et al. 2011). However, five samples collected at or near hydrograph peaks and marked with vertical dashed lines in Fig. 2a suggest that DOC concentrations sometimes increase during high flow relative to baseflow, though the broad seasonal pattern of higher concentrations during summer low flow appears stronger. This conclusion that seasonal dominance is a slightly stronger pattern than that of increases with flow during storms is confirmed by a significant ($p = 0.036$), but weak ($r^2 = 0.09$) inverse least squares linear regression relation between \log_{10} specific discharge and DOC

Table 1 Linear regression relations for FTHg concentrations with organic carbon measures for all samples, dormant season, and growing season at Fishing Brook at County Line Flow during 2007–2009

Organic carbon measure	All			Dormant season (Nov 1–Apr 30)			Growing season (May 1–Oct 31)		
	r^2	p value	Slope	r^2	p value	Slope	r^2	p value	Slope
DOC conc.	0.31	<0.001	0.17	0.42	0.003	0.24	0.25	0.013	0.22
UV ₂₅₄	0.53	<0.001	5.40	0.50	<0.001	6.57	0.57	<0.001	6.73
HPOA conc.	0.42	<0.001	0.39	0.47	0.001	0.42	0.42	0.001	0.50
HPOA %	0.21	0.002	0.079	0.14	0.11	0.077	0.49	<0.001	0.12
SUVA	0.63	<0.001	1.51	0.27	0.02	1.14	0.82	<0.001	1.57

concentrations for all samples collected at Fishing Brook; weak flow relations with DOC are a common finding in wetland-impacted streams (Hinton et al. 1997).

FTHg concentrations also were generally higher during the growing season than during the dormant season at 27 stream sites across the upper Hudson basin (Burns et al. 2012). Unlike DOC, however, FTHg concentrations increased with streamflow, a pattern observed during five different high flow periods marked with vertical dashed lines in Fig. 2a. Streamflow (\log_{10} specific discharge) was significantly ($p = 0.024$) and positively related to FTHg concentrations, consistent with a source from flushing of shallow soils as hypothesized by Schelker et al. (2011), but this relation is relatively weak ($r^2 = 0.12$), and explains little of the annual variation at Fishing Brook.

Some seasonality and response to flow conditions was observed in HPOA concentrations and SUVA₂₅₄ values, but the patterns of the two differed (Fig. 2b). SUVA₂₅₄ values sometimes followed the pattern of HPOA concentrations as in the latter half of 2008 and 2009, but sometimes show strikingly different patterns as in 2007. Overall, SUVA₂₅₄ values and HPOA concentrations are significantly ($p = 0.006$) positively correlated, but the r^2 value of 0.18 indicates a relatively weak relation. This relation improves only slightly to an r^2 value of 0.22 ($p = 0.002$) when SUVA₂₅₄ values are regressed with those of HPOA percent. Furthermore, SUVA₂₅₄ values are even less strongly correlated with those of DOC concentrations ($p = 0.10$, $r^2 = 0.06$) indicating independent variation among these two variables. In contrast, HPOA concentrations are highly correlated with those of DOC concentrations ($p < 0.001$, $r^2 = 0.92$) indicating strong seasonal synchronicity among these two quantities.

Relation of FTHg concentrations to organic carbon measures

FTHg concentrations are significantly related to all of the organic carbon measures at Fishing Brook as determined by linear regression; however, SUVA₂₅₄ values show the strongest relation with an r^2 value of 0.63 for all samples collected (Table 1; Fig. 3b). FTHg concentrations are more strongly related to SUVA₂₅₄ values than any of the other organic matter measures, including UV₂₅₄, HPOA concentration, and HPOA percent (Table 1). These results contrast with those of Dittman et al. (2009) for three northeastern U.S. streams in which FTHg concentrations showed progressively stronger relations with DOC concentrations, HPOA concentrations, and UV₂₅₄ values; r^2 values exceeded 0.90 for the latter two measures. In contrast to the strong relation between FTHg concentrations and SUVA₂₅₄ values, FTHg concentrations and DOC concentrations show an r^2 value of only 0.31 at Fishing Brook (Table 1; Fig. 3a). This finding is especially surprising because FTHg concentrations and DOC concentrations in streams and lakes with largely forested watersheds in Sweden, the Adirondacks, northeastern US, and upper Midwest typically show stronger relations with r^2 values of 0.6–0.93 (Selvendiran et al. 2008, 2009; Shanley et al. 2008; Brigham et al. 2009; Dittman et al. 2009; Eklöf et al. 2012). However, previous investigations in Fishing Brook and the upper Hudson basin noted relatively weak THg–DOC relations (Schelker et al. 2011; Burns et al. 2012). This lack of a strong Hg–DOC relation was attributed in part to the role of numerous open waters as dynamic reactors where sunlight, biological uptake, microbial processes, and interactions with lake sediment combine to alter the characteristics of DOC with respect to Hg binding and

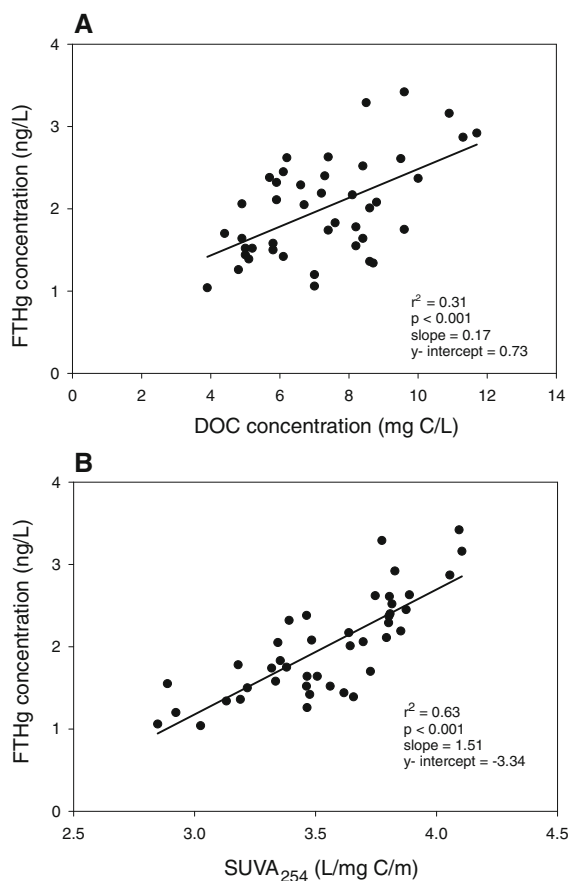


Fig. 3 FTHg concentrations and: **a** DOC concentrations, and **b** SUVA₂₅₄ values in Fishing Brook at County Line Flow for samples collected during 2007–2009. Bivariate linear regression relations are shown

to diminish THg concentrations (Schelker et al. 2011; Burns et al. 2012).

On first consideration, there is not a readily apparent reason why FTHg concentrations should be strongly related to SUVA₂₅₄ values because this organic carbon measure reflects the relative aromaticity and molecular weight of the organic matter in a sample, but is not a concentration (Weishaar et al. 2003). This study is one of several, however, in which variation in SUVA₂₅₄ values account for much of the variation in FTHg concentrations in surface waters (Table 2). Instances where the FTHg–SUVA₂₅₄ association is about as strong, or stronger than the FTHg–DOC association tend to be from synoptic-type studies in which surface waters with diverse land cover have been sampled (Hall et al. 2008; Tsui and Finlay 2011; Burns et al. 2012), with the exception of the current

study. In contrast, the FTHg–SUVA₂₅₄ relation tends to be relatively weaker than that of FTHg–DOC or FTHg–UV₂₅₄ in watersheds with more uniform, largely forested upland land cover (Schuster et al. 2008; Dittman et al. 2010). Several studies in which Hg–DOC relations were explored may not have also considered whether FTHg concentrations and SUVA₂₅₄ values were related because another measure such as DOC concentration, UV₂₅₄, or HPOA concentration was able to explain most of the variation in FTHg (Shanley et al. 2008; Dittman et al. 2009; Riscassi and Scanlon 2011; Eklöf et al. 2012). Given these results, it seems pertinent to ask why FTHg concentrations and SUVA₂₅₄ values are strongly related in many of these studies (Table 2). We hypothesize that Hg and DOC are transported to surface waters from a variety of contrasting source areas (and are also affected by sink areas such as lakes) that include riparian wetlands, forested uplands, open water and varying depths within soils. The source areas where DOC with the most aromatic character is mobilized are wetlands and in the shallowest soil horizons, which coincide with the dominant Hg mobilization zones as well. In this hypothesis, the mix of DOC and Hg from these source areas and the flow paths of water that moves these solutes to adjacent surface waters govern the apparent FTHg–SUVA₂₅₄ association. An alternative hypothesis is that the more aromatic fraction of DOC has more thiol binding sites for Hg or more binding sites in general (Ravichandran 2004; Schuster et al. 2008; Riscassi and Scanlon 2011; Tsui and Finlay 2011). We favor the first hypothesis, primarily because a large excess of reduced sulfur groups is generally available for Hg binding in most natural settings (Haitzer et al. 2002), and this excess binding ability is not expected to be greatly affected by the relative mix of aromatic and aliphatic organic matter.

The Hg–organic matter association varies by season; SUVA₂₅₄ values account for more of the variation in FTHg concentrations during the growing season ($r^2 = 0.82$, $p < 0.001$) than the dormant season ($r^2 = 0.27$, $p = 0.02$) (Fig. 4). Conversely, FTHg concentrations are more weakly associated with DOC concentrations during the growing season than during the dormant season. UV₂₅₄ values and HPOA concentrations show about the same association with FTHg concentrations during both seasons, and HPOA percent (like SUVA₂₅₄) is more strongly related with

Table 2 Summary of studies that have reported FTHg–SUVA₂₅₄ regression relations for surface waters

Reference	Study location	DOC (mgC/L)	UV ₂₅₄ (abs.)	SUVA ₂₅₄ (L/mgC/m)	Notes
This study	Fishing Brook, NY	0.31	0.53	0.63 (2.8–4.2)	One site; year round; n = 43
Burns et al. (2012)	Upper Hudson R, NY	0.32	NR	0.42 (1.6–4.5)	Summer samples; 25 sites; sampled once ea
Tsui and Finlay (2011)	MN streams	0.63	NR	0.86 (2.6–4.8)	Summer samples; 30 sites; sampled once ea; regressions for log ₁₀ FTHg
Dittman et al. (2010)	Archer Ck, NY; Hubbard Bk, NH; Sleepers R, VT	0.87	0.92	0.53 (1.9–3.9)	Sites pooled for analysis; year round; total n = 128
Hall et al. (2008)	Lower Miss. R, LA	0.69	NR	0.65 (2.6–4.5)	Lakes, rivers, and wetlands; 21 sites; sampled four times ea
Schuster et al. (2008)	Sleepers R, VT	0.31	NR	0.07 (1.3–4.0)	Snowmelt samples; ten sites; total n = 57; SUVA ₂₅₄ relation for subset of data

Values of r^2 for bivariate linear regressions of FTHg with DOC concentrations, UV₂₅₄, and SUVA₂₅₄. Approximate range of reported SUVA₂₅₄ values in parentheses

FTHg concentrations during the growing season, but the relation is not as strong ($r^2 = 0.49$, $p < 0.001$) as that of the FTHg–SUVA₂₅₄ association. A previous investigation in the upper Hudson basin also found that SUVA₂₅₄ accounted for more of the spatial variation in THg concentrations across 27 stream sites in summer than any other chemical metric (Burns et al. 2012), consistent with the results reported here.

Multiple regression models to predict FTHg concentrations

Burns et al. (2012) showed that multiple regression models that account for much of the variation in THg concentrations across the upper Hudson were stronger ($r^2 = 0.71$ – 0.84) when DOC concentrations and SUVA₂₅₄ values were included as independent variables than models obtained through either metric alone, especially in summer. Results of a forward stepwise multiple regression model for the entire period at Fishing Brook indicate that DOC concentrations and SUVA₂₅₄ values are each highly significant ($p < 0.001$, combined $R^2 = 0.75$), and indicate a modest improvement to the bivariate regression obtained with SUVA₂₅₄ values ($r^2 = 0.63$; Fig. 5a). The ΔR^2 value for DOC concentrations in the multiple regression model is 0.12, confirming that SUVA₂₅₄ accounts for most of the variation in FTHg concentrations at Fishing Brook. This regression relation

shows some bias by under-predicting at high FTHg concentrations and over-predicting at low FTHg concentrations. If log₁₀ specific discharge is added to this model ($p < 0.001$), however, R^2 increases to 0.82, and bias decreases (not shown). This model has relatively low VIF values (< 1.5) for all independent variables indicating that streamflow affects FTHg concentrations relatively independently of the other variables. SUVA₂₅₄, however, still accounts for most of the annual variation in FTHg concentrations in the model that includes streamflow.

Seasonally, SUVA₂₅₄ values and DOC concentrations are less effective at predicting FTHg concentrations during the dormant season ($R^2 = 0.55$) than the growing season ($R^2 = 0.84$) (Fig. 5b, c), though the dormant season model showed the most improvement relative to a bivariate regression with SUVA₂₅₄ values. DOC concentrations explain more variation ($\Delta R^2 = 0.42$, $p = 0.007$) in FTHg concentrations during the dormant season than do SUVA₂₅₄ values ($\Delta R^2 = 0.12$, $p = 0.05$), though both are significant dormant season predictors. This dormant season model shows greater bias than that of the annual model, underpredicting at high FTHg concentrations and overpredicting at low FTHg concentrations. In contrast, the stronger growing season model shows little bias, SUVA₂₅₄ is the dominant predictive variable ($\Delta R^2 = 0.82$), and DOC concentrations add little predictive ability ($\Delta R^2 = 0.02$, $p = 0.13$).

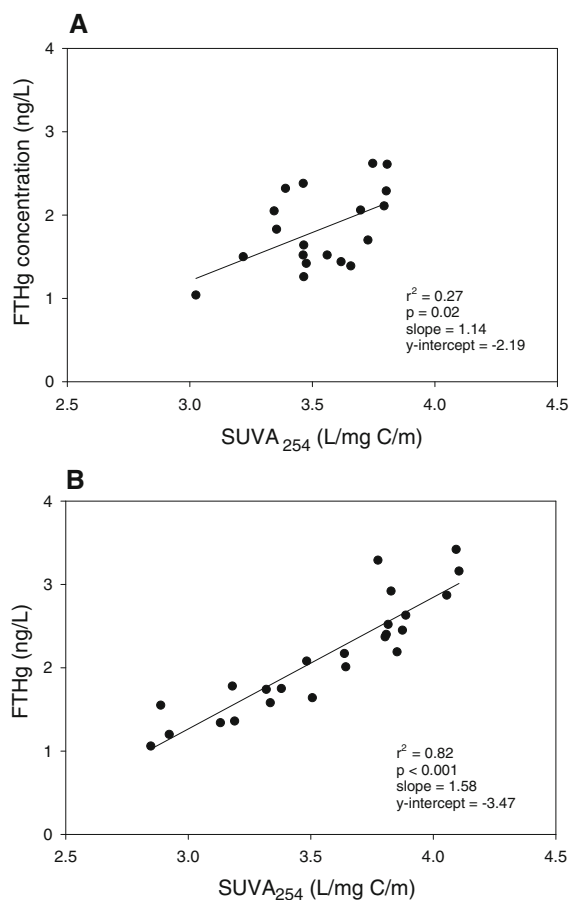


Fig. 4 FTHg concentrations and $SUVA_{254}$ values for samples collected at Fishing Brook at County Line Flow in 2007–2009 for: **a** dormant season (Nov 1–Apr 30) and **b** growing season (May 1–Oct 31). Bivariate linear regression relations are shown

The role of streamflow as an additional predictive variable in these models is quite different by season. The \log_{10} specific discharge does not improve R^2 values for the best growing season model, and has very low significance ($p = 0.93$), whereas \log_{10} specific discharge is highly significant in the best dormant season model ($p < 0.001$). This latter model has an R^2 value of 0.89, a strong improvement over the model that includes only the organic carbon measures (Fig. 5b). The low VIF value (1.1) for streamflow in the best dormant season model indicates that this variable accounts for variation in FTHg concentrations independently of the carbon measures. This finding is consistent with an hypothesis from earlier work at Fishing Brook that Hg is flushed from the shallow soil, primarily in riparian wetland areas during snowmelt and rain events, most evidently during the

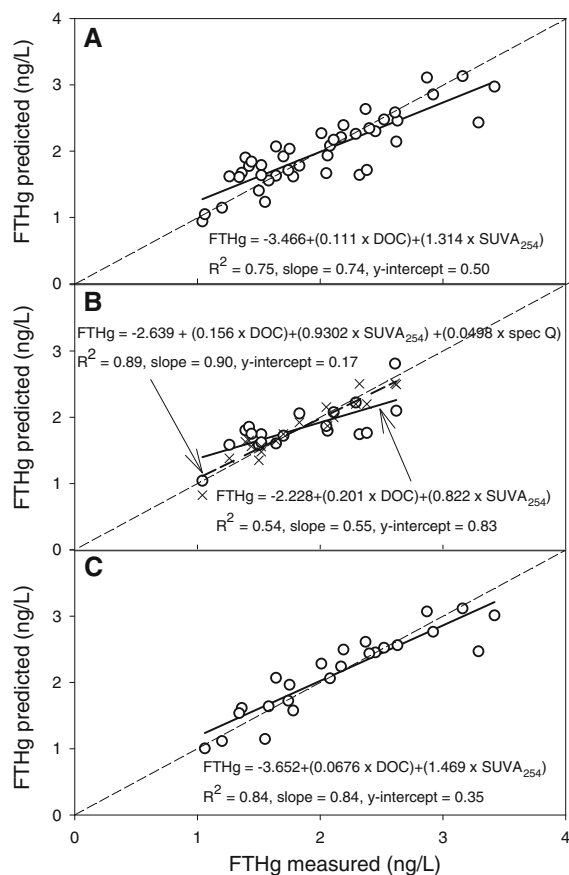


Fig. 5 Measured and predicted FTHg concentrations for Fishing Brook at County Line Flow for samples collected during 2007–2009 for: **a** entire period, **b** dormant season (Nov 1–Apr 30), and **c** growing season (May 1–Oct 31). Predicted values were derived from multiple linear regression models with DOC concentrations and $SUVA_{254}$ values as independent variables, except for an additional regression model shown in **b** with specific discharge (Q) included as an additional independent variable. Regression equations and R^2 values are shown in each panel and represented by a *solid line*, except for added model with Q shown as *long dashed line*. *Dashed lines* indicate slope of 1. Data point symbols represented by *hollow circles* except for model with Q in **b**, represented by the *times* symbol. All regression equations are highly significant ($p < 0.001$)

dormant season, and that this flushing mechanism appears to be independent of variations in DOC concentrations (Schelker et al. 2011), which are inversely and weakly related to flow as stated previously. In contrast, variation in growing season FTHg concentrations in Fishing Brook is strongly associated with variation in $SUVA_{254}$ values, with little influence from either DOC concentrations or flow. This finding raises important questions as to the source areas and

processes that affect variation in $SUVA_{254}$ values in this basin, which are explored further in the next section.

Relation of $SUVA_{254}$ values to landscape metrics

Given the apparent importance of $SUVA_{254}$ values in explaining growing season variations in FTHg concentrations at Fishing Brook, the role of several landscape metrics are explored here to gain insight to the sources of variation in $SUVA_{254}$ values across the upper Hudson basin. Multiple regression models that best account for variation in $SUVA_{254}$ values were developed for synoptic samples collected at 27 stream sites across the upper Hudson River basin during 2006 and 2008 (Fig. 1). These sites range from steep upland watersheds to lowland streams with abundant riparian wetlands and varying influence from numerous open water bodies in the basin and are described in greater detail by Burns et al. (2012). Several of these sampling sites are within the Fishing Brook watershed discussed previously.

$SUVA_{254}$ values for synoptic samples collected in spring are well explained ($R^2 = 0.89$) by a model that includes slope, percent riparian area (ripct), and an OWI (Fig. 6a). $SUVA_{254}$ values are highly correlated ($p < 0.001$) in bivariate regressions with slope (inversely), and percent riparian area; $r^2 = 0.82$ and 0.70 , respectively. In contrast, $SUVA_{254}$ values are not strongly correlated ($p = 0.23$) with OWI in a bivariate regression, but this metric is significant ($p = 0.02$, $\Delta R^2 = 0.04$) in the best multiple regression model indicating a minor role for open water. These spring-time results suggest that higher $SUVA_{254}$ values originate from soils in areas with gentler slopes, and in riparian areas that are commonly occupied by wetlands (percent wetland area strongly correlated ($r^2 = 0.66$) with $SUVA_{254}$), and a secondary role for open waters as a source of lower $SUVA_{254}$ values. These results are consistent with those of previous investigations in Oregon and in Sweden that have found higher $SUVA_{254}$ values associated with riparian/wetland areas than values derived from forested upland areas (Hood et al. 2006; Ågren et al. 2008).

The multiple regression modeling results for synoptic samples collected in summer show some differences with those collected in spring (Fig. 6b). The best model in summer has an R^2 of only 0.49, shows some bias (high at low $SUVA_{254}$, low at high $SUVA_{254}$), and indicates

that about half the variation in $SUVA_{254}$ values cannot be accounted for by these landscape metrics. This model includes slope and OWI as in the spring model, but also includes TWI and the small open water metric. Both accumulated area (analogous to drainage area) and slope are part of the TWI calculation, and reflect the tendency for a higher water table as values increase (Beven and Kirkby 1979). The TWI is inversely related ($r^2 = 0.19$, $p = 0.02$) to slope in the upper Hudson basin, but is more strongly (and positively) related to the percent open water in each sub-basin ($r^2 = 0.27$, $p = 0.006$), indicating that basins with high TWI values have gentler slopes, but also more open water. These two influencing factors are expected to have opposing effects on $SUVA_{254}$ values.

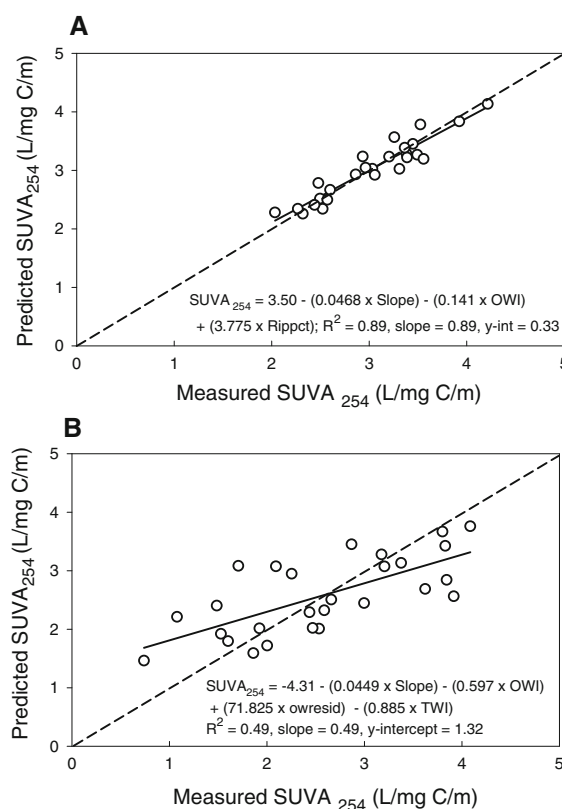


Fig. 6 Measured and predicted $SUVA_{254}$ values for synoptic samples collected during 2006 and 2008 at 27 surface water sites in the Upper Hudson River basin: **a** spring (May), and **b** summer (July and Aug). Predicted values were derived from multiple linear regression models that considered several landscape metrics as discussed in text. Regression equations and R^2 values are shown in each panel and represented by a solid line. Dashed lines indicate slope of 1. Regression in **a** is highly significant, $p < 0.001$, regression in **b** is significant, $p = 0.004$

Overall, three of four independent variables in the multiple regression model for $SUVA_{254}$ based on summer synoptic samples are related in part to open water influence, and suggest stronger influence of open water on $SUVA_{254}$ values in summer than in spring. The inverse relations with OWI and TWI indicate open water is associated with lower $SUVA_{254}$ values. This finding suggests that low $SUVA_{254}$ values may be derived in part from autochthonous processes such as decomposition of algae in the water column (Zumstein and Buffle 1989; Rostan and Cellot 1995). Photodegradation of allochthonous DOC may also contribute to declining $SUVA_{254}$ values associated with open water influence (Minor and Stephens 2008). Regardless of the processes responsible, $SUVA_{254}$ values often reach a minimum and the influence of autochthonous DOC reaches a maximum during summer low flow, particularly in lake-influenced systems (Zumstein and Buffle 1989; Hood et al. 2005) when sunlight and water temperature are at seasonal maximum values, consistent with the results presented here for summer synoptic samples. In contrast, the positive relation of $SUVA_{254}$ values with the small open water variable is consistent with observations that these open waters are commonly beaver ponds or transitional wetlands in this basin, and thus behave more similarly to wetlands as a source of Hg (Driscoll et al. 1998) and higher $SUVA_{254}$ values.

Although synoptic samples were generally collected during similar low-to-moderate flow conditions, flows were greater in spring (mean = 1.8 mm/day, RSD = 55 %) than in summer (mean = 1.3 mm/day, RSD = 109 %). Because DOC and Hg concentrations are often strongly related to streamflow and the latter may be similarly strongly related to $SUVA_{254}$ values (Hood et al. 2006; Schuster et al. 2008; Dittman et al. 2010), specific discharge was considered as a potential additional explanatory variable in the best multiple regression $SUVA_{254}$ models. Consideration of streamflow is further reinforced by a significant bivariate regression relation ($r^2 = 0.32$, $p = 0.002$) between $SUVA_{254}$ values and specific discharge for samples collected during the growing season at Fishing Brook. However, specific discharge was not significant ($p > 0.05$) in either spring or summer multiple regression synoptic models. Thus, inclusion of flow did not improve model predictions from those obtained with just landscape metrics. This analysis has limitations, however, because these synoptic samples do not well

represent high flow conditions, nor is the role of storms reflected by the samples.

$SUVA_{254}$ values are significantly inversely correlated with streamflow for samples collected in spring ($r^2 = 0.36$, $p = 0.002$), and positively correlated for samples collected in summer ($r^2 = 0.26$, $p = 0.006$). This difference may reflect the wetter conditions and higher water tables in this basin during spring than summer (Schelker et al. 2011). A previous study in Sweden in a similar heterogeneous landscape of upland forests and lowland riparian wetlands concluded that runoff in spring from forested upland areas (with lower $SUVA_{254}$ values than in riparian areas) has a greater influence on stream DOC, and $SUVA_{254}$ accordingly decreased with flow (Ågren et al. 2008). During summer, in contrast, when riparian saturation deficits are greatest, riparian influence increased with flow, and $SUVA_{254}$ values accordingly increased with flow.

There are clearly sources of variation in $SUVA_{254}$ values not fully captured by models derived from the landscape metrics explored here, as evidenced by the R^2 value of 0.49 for summer samples. Carbon chemistry is complex with multiple molecular structures identified in DOC (Leenheer and Croué 2003), but the success of the modeling approach employed here, particularly for the spring synoptic samples, points to the principal sources of variation in $SUVA_{254}$ as it represents the relative aromaticity of DOC and strong association with Hg. A broad pattern of $SUVA_{254}$ values across the 27 sub-basins is observed when these data are divided into quartiles by percent riparian area and compared with groundwater seep samples collected in the largest riparian wetland area in the Fishing Brook basin (Fig. 7a). An analysis of variance on ranks (because data did not pass test for normality) indicated a statistically significant difference ($p < 0.001$) among these groups. Pairwise comparisons, however, showed that not all quartiles are statistically distinct, but that $SUVA_{254}$ values generally increase as riparian area increases. The highest and most distinct values are in the uppermost quartile and in the groundwater seeps, the latter of which may be viewed as a “pure” riparian wetland source of DOC. The range in median $SUVA_{254}$ values across the quartiles and seeps is about 1.5 L/mg C/m.

Open water influence on $SUVA_{254}$ values can be similarly observed by examining the synoptic data by quartiles of the OWI (Fig. 7b). An analysis of variance

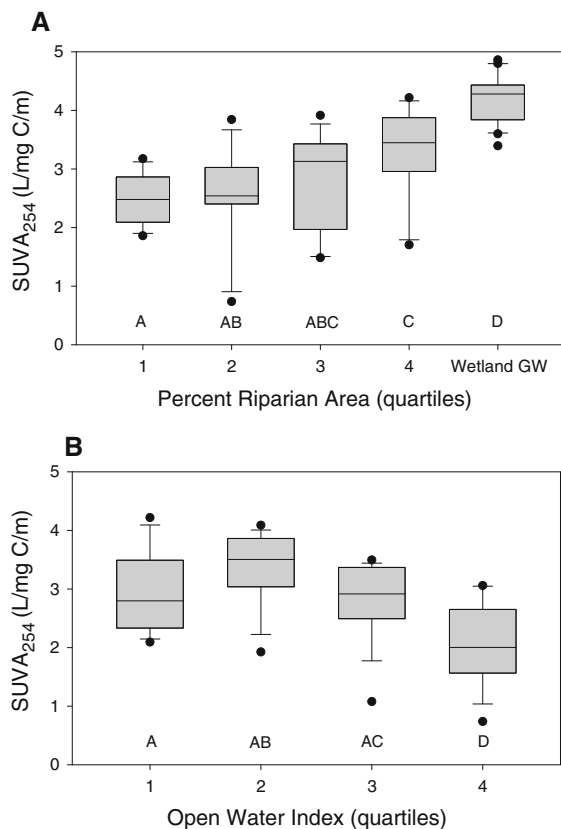


Fig. 7 Box plots of data from Upper Hudson River basin: **a** $SUVA_{254}$ values by percent riparian area quartiles for synoptic samples collected during 2006 and 2008 at 27 surface water sites and in wetland groundwater seeps collected during 2007–2009, and **b** $SUVA_{254}$ values by OWI quartiles for same synoptic samples shown in **a**. Capital letter designations below each box indicate statistical similarity and differences among pairwise comparisons of each quartile or group

indicated a statistically significant difference among these quartiles. A general decrease in $SUVA_{254}$ values is observed across the quartile range, with the exception of the first to second quartiles. The highest quartile shows the lowest and statistically most distinct $SUVA_{254}$ values (based on pairwise comparisons) similar to that observed for the highest percent riparian area quartile. Also, the range in median values among the OWI quartiles is slightly greater than that of the percent riparian area quartiles.

Relevance of results to methyl Hg transport

Although the analysis above focused on FTHg, filtered methyl Hg (FMHg) concentrations were also measured in the larger study from which this paper is

derived, and these data have been discussed in previous papers (Bradley et al. 2011; Schelker et al. 2011; Burns et al. 2012). Broadly, FMHg concentrations are significantly related ($p < 0.05$) with FTHg concentrations and with $SUVA_{254}$ values. These relations, however, do not have strong explanatory ability ($r^2 = 0.1$ – 0.2), reflecting the influence of different processes that control the production of FMHg and FTHg as well as varying dependence on organic carbon fractions for transport of these Hg species to surface waters. Nonetheless, the discussion of FTHg transport is believed to have some broad transfer value to understanding FMHg transport as well because FTHg may be later transformed to FMHg.

Conclusions

The results of this study indicate an important role for $SUVA_{254}$ as a predictor of temporal and spatial variation in FTHg concentrations in the upper Hudson River basin. At Fishing Brook, $SUVA_{254}$ values explained twice the annual variation of FTHg concentrations than could be explained by DOC concentrations. Multiple regression models that included both variables accounted for an even greater amount of this variation in FTHg concentrations for all samples collected at Fishing Brook, but especially during the Nov–Apr dormant season. $SUVA_{254}$ alone, however, was a notably stronger predictor of FTHg concentrations in the May–Oct growing season. This seasonal difference is believed to result largely from the opposing influence of (1) gentler slopes and more extensive riparian area as a source of higher $SUVA_{254}$ values and FTHg concentrations, and (2) open water as a source (with the exception of small open water bodies) of lower $SUVA_{254}$ values and FTHg concentrations. These data are also consistent with an additional role for increasing contributions of runoff originating from shallow flow paths during events associated with greater $SUVA_{254}$ values and FTHg concentrations; however, the relationships of $SUVA_{254}$, DOC, and FTHg with streamflow are relatively weak, suggesting that flow paths alone are only a minor explanatory variable (though samples collected in this study may not have adequately characterized stormflow). We believe these results generally reflect the manner in which the seasonal

dynamics of upland versus riparian runoff sources combine with open water influence and flow-related variation to produce an apparent FTHg–SUVA relation. The aromaticity of DOC associated with higher SUVA₂₅₄ values may contribute to the patterns observed here, but is probably not the most important factor because of the excess of strong Hg-binding sites generally available on DOC regardless of its relative aliphatic versus aromatic character. We believe these findings are especially pertinent to larger heterogeneous basins that encompass uplands, riparian wetlands, and open water bodies such as those commonly found in the northeastern US, southeastern Canada, and the upper Midwestern US (Lehner and Döll 2004), but these results await further confirmation through additional studies in large, heterogeneous basins.

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